Corrigendum


Todd C. Mockler, Simon Chan, Ambika Sundaresan, Huaming Chen, Steven E. Jacobsen, Joseph R. Ecker

*Plant Biology Laboratory, The Salk Institute for Biological Studies, La Jolla, CA 92037, USA
bGenomic Analysis Laboratory, The Salk Institute for Biological Studies, La Jolla, CA 92037, USA
cDepartment of Molecular, Cell and Developmental Biology, University of California, Los Angeles, CA 90095

Simon Chan, Ambika Sundaresan, Huaming Chen, and Steven E. Jacobsen should have been listed as authors, and Huaming Chen’s name therefore should not have appeared in the Acknowledgments section.

The correct author line appears above.
Applications of DNA tiling arrays for whole-genome analysis

Todd C. Mockler\textsuperscript{a}, Joseph R. Ecker\textsuperscript{a,b,*}

\textsuperscript{a}Plant Biology Laboratory, The Salk Institute for Biological Studies, La Jolla, CA 92037, USA
\textsuperscript{b}Genomic Analysis Laboratory, The Salk Institute for Biological Studies, La Jolla, CA 92037, USA

Received 7 July 2004; accepted 14 October 2004

Abstract

DNA microarrays are a well-established technology for measuring gene expression levels. Microarrays designed for this purpose use relatively few probes for each gene and are biased toward known and predicted gene structures. Recently, high-density oligonucleotide-based whole-genome microarrays have emerged as a preferred platform for genomic analysis beyond simple gene expression profiling. Potential uses for such whole-genome arrays include empirical annotation of the transcriptome, chromatin-immunoprecipitation-chip studies, analysis of alternative splicing, characterization of the methylome (the methylation state of the genome), polymorphism discovery and genotyping, comparative genome hybridization, and genome resequencing. Here we review different whole-genome microarray designs and applications of this technology to obtain a wide variety of genomic scale information.

© 2004 Elsevier Inc. All rights reserved.

Contents

Applications of whole-genome arrays (WGAs) ............................................. 0

Unbiased measure of transcriptional activity ............................................. 0

Alternative splicing ................................................................. 0

Identification of RNA-binding protein targets ............................................ 0

Genome analysis............................................................... 0

ChIP-chip ................................................................. 0

Methylome analysis ............................................................... 0

Polymorphism analysis and genome resequencing ........................................ 0

Tiling array design considerations .................................................. 0

Challenges and future directions ..................................................... 0

Acknowledgments ................................................................ 0

References ................................................................ 0

The completion of numerous genome sequences has introduced an era of whole-genome study. Gaining a more complete understanding of the genome’s information content will dramatically improve our understanding of various biological processes. In parallel with the sequencing of entire genomes, recent advances in microarray technologies have made it feasible to interrogate an entire genome sequence with arrays. Such high-density whole-genome DNA microarrays can be used as a generic platform for numerous experimental approaches to decode the information contained within the genome. In this review, we discuss several approaches using high-density whole-genome oligonucleotide microarrays for transcriptome characterization, novel gene discovery, analysis of alternative splicing, mapping of regulatory DNA motifs using the chromatin-immunoprecipitation (ChIP) chip, whole-genome DNA methylation analysis, polymorphism analysis, comparative genome analysis, and genome resequencing (Fig. 1).
There are two general types of high-density microarray platforms that are most widely used. High-density oligonucleotide arrays contain relatively short (~100-mer) probes synthesized directly on the surface of the arrays by photolithography using light-sensitive synthetic chemistry and photolithographic masks [1–3], an ink-jet device [4], or programmable optical mirrors [5–7]. Oligonucleotide arrays can be made with >6,000,000 discrete features per chip, with each feature comprising millions of copies of a distinct probe sequence. The second array platform is made by mechanically printing/spotting probes, generally amplified PCR products, oligonucleotides, or cloned DNA fragments, onto glass slides. This type of array generally has a much lower feature density than the in situ synthesized oligonucleotide arrays, typically of about 10,000–40,000 spots per chip.

Because oligonucleotide arrays offer a much higher feature density and high reproducibility, and probes can be synthesized to represent virtually any sequence of a finished genome, they are the preferred platform for whole-genome analysis. Moreover, the relatively short probe length combined with the flexibility of using multiple overlapping probes representing the same genomic region makes oligonucleotide arrays ideal for detecting the broadest range of genomic features, including small polymorphisms and splice variants, and the specificity also potentially allows repetitive regions or gene family members to be distinguished.

For the purpose of this review, we will distinguish whole-genome tiling arrays designed to interrogate an entire genome in an unbiased fashion [8–11] from quasi-whole-genome (nontiling) arrays, or whole-genome expression arrays, that represent the known and predicted (annotated) features of a genome, such as exons or splice junctions on a whole-genome scale [12,13]. Nonoverlapping or partially overlapping probes (Fig. 2A) may be tiled to cover the entire genome end to end, or the probes may be spaced at regular intervals (Fig. 2B). Such an unbiased approach allows researchers to analyze various features of the genome, including evidence of transcriptional activity, binding of transcriptional regulators, and DNA methylation, at high resolution without reference to prior annotations. Other array designs rely on prior genome annotation to interrogate a particular subset of features of an entire genome (Fig. 2C). These arrays are clearly limited by the quality and completeness of the annotations on which they are based.

Applications of whole-genome arrays (WGAs)

Unbiased measure of transcriptional activity

It is well known that having a finished genome sequence is not sufficient to identify all of the transcription units, as computational gene prediction methodologies are fraught with errors. While traditional molecular approaches to identifying genes, including cloning and sequencing large collections of cDNAs, have succeeded at identifying expressed transcripts for tens of thousands of genes [11,14–17], they eventually reach a point of greatly diminished returns. Transcripts that are low abundance or expressed in rare cell types or in response to specific stimuli may never be identified by these methods. Microarrays can be used to circumvent some of these problems, allowing confirmation of the predicted gene models as well as being a tool for new gene discovery. One study [13] used ink-jet-fabricated oligonucleotide arrays to study gene transcription and transcript structure on human chromosome 22 and the human genome as a whole. Exon-scanning arrays containing relatively long (50- to 60-mer) oligonucleotide probes were hybridized with labeled cDNAs derived from various cell lines and tissues and the resulting probe intensities were used to identify expressed exons, model gene structures, and compare differential expression across conditions. Overall, expression was detected for ~57% of Genscan-predicted genes; however, this study did not interrogate most of the nonrepetitive sequence on chromosome 22. Because the exon-scanning arrays were designed using only known and computationally predicted exons, they were of limited use for discovering novel genes or gene features, such as terminal
exons that are often missed by the gene prediction algorithms. For some genomic regions, tiling arrays with partially overlapping (10-base increments) 60-mer probes were used to demonstrate the utility of high-resolution tiling arrays for refining and confirming gene structures predicted by the exon arrays. These approaches confirmed the utility of WGAs for at least a first stage of empirical validation of genome annotations. However, none of the arrays were able to provide an unbiased view of the genome and thus could not interrogate portions of the genome outside of the annotations.

Until recently, the traditional view of the genome was that “genes” were the regions of the genome that encoded proteins or structural RNAs; the regions upstream of genes were considered regulatory sequences (promoters), and much of the rest was considered “junk.” Recent analyses of the transcriptome using unbiased WGAs have challenged this view. Studies involving arrays containing PCR fragments covering all unique regions of human chromosome 22 [18] or high-density oligonucleotide tiling arrays covering essentially all nonrepetitive regions of human chromosomes 21 and 22 [8–10] or the entire Arabidopsis genome [11] have been used to map the sites of transcriptional activity in an unbiased fashion. These studies revealed that up to 10 times more expression than predicted from prior EST or cDNA evidence or gene predictions was evident in the human and plant genomes, although questions remain as to whether these observations reflect so-called “transcriptional noise.”

High-density oligonucleotide tiling arrays have been used to interrogate chromosomes 21 and 22 via 25-mer probes spaced on average every 35 bp [8–10]. The arrays were probed with labeled targets prepared from poly(A)⁺ mRNAs isolated from several different human cell line and tissue samples. The results indicated that a much larger portion of the human genome is transcribed than was previously appreciated, with as much as 90% of the observed transcription occurring outside of known or otherwise annotated exons. Using a completely different microarray platform comprising PCR fragments representing most of the non-repetitive sequence of human chromosome 22, Rinn et al. [18] found at least twice as much transcribed sequence as expected based on the annotation. Rinn et al. also compared the expressed sequences of chromosome 22 predicted by their study with those predicted by Kapranov et al. [9] and found extensive overlap, with ~90% of the ~2500 hybridizing PCR fragments on their array being detected as expressed in Kapranov’s study. Thus considering that at least half of the observed transcription in these studies that used different microarray platforms was outside of any known annotation, ~45% of the novel transcription was observed in both studies. The hypothesis that these novel transcripts derived from previously unannotated portions of the genome possess some biological function instead of merely being transcriptional noise is supported by evidence of evolutionary sequence conservation for some ~20 [10] to ~44% [18] of these transcripts based on comparisons to the mouse genome. Many of the novel transcription units observed in these studies correspond to so-called noncoding RNAs or untranslated (UTRNA) transcripts, which may be at least as prevalent as transcripts encoding proteins. Less than a quarter of the transcription units found to occur outside of annotated gene models appear to contain ORFs longer than 75 bp, suggesting that most of the novel transcripts identified...
may represent noncoding RNAs \[8,10\]. Further studies are needed to confirm these striking findings and to elucidate the functional relevance of these novel transcription units. Consistent with these results, a computational analysis of the mouse transcriptome using full-length cDNA sequences revealed that about ~73\% of novel (unannotated) mouse transcripts were noncoding \[15\]. Some of the novel transcripts predicted using the tiling arrays were further characterized by RT-PCR and Northern blot analysis and found to be spliced and/or polyadenylated. These utRNAs share many of the properties of coding transcripts and are likely to have important functions in the cell, thus challenging the basic tenet of the central dogma of molecular biology, namely that the information content of the genome flows from DNA to RNA to protein \[19\].

WGAs have also been useful for studying another poorly understood aspect of the transcriptome, natural antisense transcripts (NATs) \[20–22\], because they can simultaneously monitor gene expression on both strands of a genome. In the oligonucleotide tiling array studies of human chromosomes 21 and 22, a significant proportion of exonic sequence was found to exhibit antisense transcription \[9,10\]. Unlike the prior study \[9\] that used non-strand-specific double-stranded cDNA made with random-primed reverse transcription of cytosolic poly(A)+ RNA samples, Kampa et al. \[10\] hybridized end-labeled RNA targets from two different cell lines to the chromosome 21 and 22 arrays. About 11\% of the observed transcription in terms of base pairs of known exons, mRNAs, and ESTs was found to be antisense. About half of the observed novel intronic transcription was antisense to well-characterized introns, consistent with the study by Rinn et al. \[18\] that showed that about 50\% of transcription within annotated introns was antisense. Altogether at least 20\% of the total base pairs of chromosomes 21 and 22 exhibited antisense transcription, consistent with other recent reports of large-scale antisense transcription in eukaryotes \[21,23–28\].

The first eukaryotic genome to be entirely represented on arrays was the Arabidopsis genome. Tiling arrays representing the entire genome sequence with 25-mer probes tiled end to end (i.e., 25-bp resolution) were used to profile labeled cRNA targets derived from different tissues and/or developmental stages of Arabidopsis plants \[11\]. Numerous intergenic regions containing no annotated gene models were found to be transcribed (Fig. 3A), computational gene models were confirmed or corrected (Figs. 3B and 3C), and untranslated regions (5‘ and 3‘ UTRs) of known genes were extended (Fig. 3D). Novel transcripts were also identified within the genetically defined centromeres. Finally, and most unexpectedly, a strikingly large amount (>30\%) of antisense transcriptional activity was observed from known genes with sense strand expression (i.e., those with ESTs or cloned full-length cDNAs) \[11\] (Mockler and Ecker, unpublished). Taken together with the NAT predictions based on tiling array studies \[10,18\], these observations point to a significant amount of NAT expression in higher eukaryotes that cannot be readily explained as being an artifact of the labeled target/probe preparation because several of these studies involved hybridization of strand-specific labeled targets \[10,11\] or probes \[18\] to microarrays. Moreover these findings are consistent with NAT predictions based on transcript sequence evidence \[23–25,27\], although the functional significance of this phenomenon remains largely unknown.

Recently WGAs were also used to study gene expression patterns in the malaria parasite Plasmodium falciparum \[29\]. Nine different stages of the malaria parasite’s life cycle in both the mosquito and the human host were profiled using oligonucleotide arrays containing probes representing both strands of the predicted coding sequences, including both the mitochondrial and the plastid genomes, and noncoding sequences. On average probes were spaced in about 150-base intervals.

Collectively, these studies show that tiling arrays are a powerful tool for characterizing the transcriptome. However, it should be noted that in several of these studies a portion of the data was generated from transcript populations derived from cell lines that are likely to have gross genomic DNA alterations. Thus, some of the observed transcriptional activity may reflect these genomic differences from normal wild-type cells. Nevertheless, these recent discoveries made using tiling arrays signal a paradigm shift in how we think about the transcriptome and highlight some limitations of relying on cDNA and EST sequencing projects for gene discovery. Given the way cDNA libraries are generally constructed and queried, it seems likely that specific classes of transcripts [for example, extremely low abundance, very short, or atypical (non-poly(A)+, noncoding, those with extensive secondary structure, etc.)] may never be identified and sequenced by traditional approaches. Regardless of which type of tiling array platform is used, the hybridization signals and corresponding gene structures remain predictions that must be confirmed by RT-PCR amplification, cloning of full-length cDNAs, and sequencing. An alternative approach that may prove more efficient would be first to attempt to establish the functional relevance of novel transcripts identified using tiling arrays. For example, patterns of coexpression/coregulation between the novel predicted transcripts and the known transcripts could be scored and used to prioritize the predicted transcripts for further characterization. Empirical approaches complementary to WGA analysis will result in significantly improved, experimentally validated genomic annotations and enable the generation of previously identified cDNA clones that are valuable resources for other experiments. However, these validation efforts remain laborious and time-consuming.

**Alternative splicing**

Another use of WGAs has been for the discovery and characterization of differentially/alternatively spliced tran-
Fig. 3. Comparison of *Arabidopsis* transcription units using computational and WGA analyses. (A) Example of a novel unannotated gene transcription in an intergenic region, (B) a computationally derived gene model (At3g14670) that is annotated on the wrong strand, (C) gene model confirmation and correction for an annotated hypothetical gene (At2g04135), and (D) a novel 5’ exon for a known gene (At5g42190) from *Arabidopsis* whole-genome tiling array data visualized using the ChipViewer software (H. Chen and J. Ecker, unpublished and http://signal.salk.edu/cgi-bin/atta).
scripts. Alternative splicing is an important mechanism of genetic regulation in higher eukaryotes that increases protein diversity by allowing multiple functionally distinct proteins to be encoded by a single gene, with some genes potentially generating thousands of isoforms [30–33]. Some RNA-binding proteins act as trans-acting splicing factors that bind to numerous exonic splicing enhancers within transcripts to promote inclusion of alternative exons with weak splice sites in the pre-mRNA, while others bind to intronic splicing modifier sequences. Alternative splicing is hypothesized to result from a dynamic antagonism between these trans-acting factors that bind to regulatory elements in the pre-mRNA [30,34]. Numerous computational studies have examined alternative splicing by analysis of EST and cDNA transcript sequence evidence [35–38]. Although these computational approaches have identified thousands of alternatively spliced transcripts, they are constrained by the amount and quality of the available transcript sequence data. Thus human or mouse, with about 10 times more ESTs than other organisms (http://www.ncbi.nlm.nih.gov/dbEST/), may seem to have more extensive alternative splicing due to simply having more EST sequences available. WGA{s} potentially offer a powerful platform for both identifying new alternatively spliced transcript forms and characterizing the splicing of known transcripts in diverse collections of samples such as different cell types, tissues, developmental stages, or drug treatments. In a pioneering study demonstrating the utility of microarrays for studying splicing, microarrays containing oligonucleotide probes corresponding to splice junctions, introns, and exons were used to characterize alternative splicing for all intron-containing genes of yeast [39]. More recently, ink-jet-synthesized arrays have been designed that specifically examine alternative splicing of known gene models [12,40]. These arrays were not whole-genome tiling arrays per se and thus have an inherent limitation of being biased for known genes and predicted gene models. In one study, Castle et al. [40] demonstrated the utility of ink-jet oligonucleotide arrays for analyzing transcript structure and alternative splicing. Optimal probe lengths, tiling densities, probe overlap requirements, and hybridization conditions were assessed for the detection of various transcript features, including differentiating exons from introns, calling exon edges, and monitoring splice junctions. These pilot experiments determined that the optimal design for identifying exons used 50- to 60-mer probes, exon edges could be best defined using 35- to 40-mer overlapping probes tiled across splice junctions, and 30- to 40-mer splice-junction probes centered exactly on the splice junction or slightly offset were optimal.

In a follow-up study, ink-jet-fabricated splice-junction arrays were used for a whole-genome survey of alternatively spliced transcripts [12,40]. A set of five ink-jet arrays comprising about 125,000 36-mer splice-junction probes was used to monitor the splicing of ~10,000 human genes. Labeled cDNA targets from 52 tissue and cell lines were hybridized to these arrays in duplicate. Half of the alternative splicing predictions based on the array data could be validated by RT-PCR and sequencing. This approach has provided strong evidence for alternative splicing in thousands of genes and identified alternative splice variants not previously predicted from cDNA evidence. Also, completely novel alternative splicing was found for genes not previously known to be alternatively spliced. However, the array-based approach to analyzing alternative splicing also has limitations. For example, for relatively rare splicing events, a large number of distinct tissues or cell populations must be surveyed to obtain sufficient confidence to call a splice variant at a specific splice junction. In addition, the splice junction arrays cannot distinguish whether two splicing events observed in one sample occur in the same or distinct transcripts, if the two isoforms are expressed at similar levels. Moreover, if two splicing events are mutually exclusive, then prior knowledge is required to design probes capable of recognizing both forms. An additional challenge is that splice-junction probes are constrained by their positions in the transcripts and therefore cannot be chosen for optimal binding characteristics (probe affinity, secondary structure, and specificity).

Unbiased WGAs have also been used to monitor one type of alternative splicing event—exon-skipping [9,10]. By comparing the expression levels of all of the exons within genes, it was shown that a large fraction (~80%) of expressed genes on chromosomes 21 and 22 exhibit exon skipping. This estimate is significantly higher than previous predictions [36,41] based on computational analysis of ESTs and cDNAs. However, it should be noted that estimates of alternative splicing based on expressed sequence data (ESTs and cDNAs) can be problematic for a few reasons and thus may be underestimates. First, the EST and cDNA data are biased against low-abundance transcripts that may not be well represented in the sequence databases, whereas the array approach can detect alternative splicing even for low-abundance tissue-specific transcripts. In addition, alternative splicing predictions based on sequence data are likely biased toward the termini of transcripts due to the preponderance of end-sequence reads among ESTs and oligo(dT)-based priming for reverse transcription.

Much work remains to be done to make microarrays a preferred platform for routine analysis of alternative splicing, and it is not yet clear which whole-genome array design is best for this application. Should unbiased tiling WGAs, splice-junction arrays, exon scanning, or some combination of all of the above be used to study alternative splicing? Regardless of the general array design chosen to study splicing, a reasonable approach would be to integrate all available prior evidence for alternative splicing (such as EST- and cDNA-based predictions) into both the array design and the probe selection as well as the analysis of the resulting data to increase the confidence of the microarray-based alternative splicing calls. Many other equally important questions remain as well. For splice-junction analysis,
what is the best density of the junction probes, a single junction oligo or multiple tiled oligos? How many probes per exon are optimal for exon-scanning designs? Should a unbiased approach be taken, or should the probes be chosen based on known or predicted transcript structures? One advantage of knowing multiple isoforms for a given gene beforehand is that specific probes for each of the different isoforms can be included on the array. Splice-junction probes can distinguish different isoforms present in one sample as a mixture and simultaneously assess the relative expression levels of the different isoforms in the mixture. In the case of an isoform mixture, it is likely that tiling array probe signals would reflect a merged or composite structure of the transcript. One distinct advantage of interrogating splice junctions is that because the junction probes tile across nonadjacent genomic sequences (i.e., the spliced exons), they can indicate which genomic regions, the so-called “trans-frags,” are actually connected together in a transcription unit. This information cannot be readily inferred from whole-genome tiling arrays without further analyses, such as carrying out 5’ and 3’ RACE followed by hybridization of the RACE products onto tiling arrays or by doing directed RT-PCR. Regardless of the array design used to study alternative splicing, the potentially new splice forms that are predicted using these arrays must be validated by further experimentation such as sequencing of RT-PCR products. A combined approach would use an unbiased whole-genome tiling array to screen for new alternative splice forms. These candidates could then be verified with arrays specifically designed to verify the splice junctions.

Identification of RNA-binding protein targets

Eukaryotic genomes generally encode hundreds of predicted RNA-binding proteins (RBPs) that are likely involved in various aspects of posttranscriptional gene regulation, including transcript splicing, RNA stability, transcript localization, and translational regulation [42,43], yet the transcript targets have been comprehensively identified for relatively few RBPs [44–50]. Another potential application of WGas is for the systematic identification of the transcript targets of RBPs following the purification of the RBP-bound RNAs either using an affinity-purification approach [45] or following immunoprecipitation with antibodies directed against the RBP of interest [49,51].

In a recent study, DNA microarrays were used to identify the specific RNA targets of five affinity-tagged *Saccharomyces cerevisiae* Pumilio–Fem-3-binding factor (Puf) family RBPs [45]. TAP-tagged Puf proteins were purified from whole-cell yeast extracts and the mRNAs that interacted with the tagged Puf proteins were isolated. Then, labeled cDNAs were prepared from these transcripts and hybridized to DNA microarrays containing all of the known and predicted nuclear yeast ORFs and introns and the mitochondrial genome. As a control for nonspecific binding of RNAs, the procedure was performed in parallel using cells lacking the TAP-tagged Puf proteins. Transcripts representing more than 700 distinct ORFs (~12% of yeast genes) were identified as interacting with at least one Puf protein, and 90 transcripts were found to interact with more than one Puf protein. The specific interactions with some Puf targets were confirmed using a yeast three-hybrid assay [45,52]. Each of the five Puf proteins interacted with specific sets of mRNAs that could be classified by their functional annotations and by subcellular localization. For three of the Puf proteins studied, distinct sequence elements identified in the 3’ UTRs were implicated in specific interactions between the Puf proteins and their targets.

Although expression arrays alone can be used to identify RBP target genes for many of the known and predicted protein coding genes, tiling arrays may allow the association of specific interactions of RBPs with certain alternatively spliced transcript variants. Such an approach could have important implications for understanding posttranscriptional regulation of alternatively spliced transcripts, including their localization, stability, and translation. In addition, tiling arrays may allow identification of RBP targets among the plethora of recently discovered trans-frags that lie in the unannotated intergenic regions as well as antisense RNAs [8–11,53]. A complementary approach might involve comparing transcript populations from RBP mutants or cells overexpressing an RBP using splicing or tiling arrays to identify transcript targets of specific RBPs.

Genome analysis

ChIP-chip

In addition to simply characterizing the transcribed portions of the genome, another goal of genomic studies is the identification of the complement of regulatory DNA sequences that are bound by transcriptional regulators. Chromatin immunoprecipitation is a method that has been developed to isolate and identify the in vivo DNA sequences that are bound by a transcription factor [54]. Briefly, crosslinked chromatin (DNA/protein) complexes are extracted from a tissue of interest and sheared, typically by sonication, down to relatively short (<1 kb) fragments. The chromatin fragments containing the transcription factor of interest are then immunoprecipitated using an antibody against the transcription factor (or a fused epitope tag) and the target sequences are subsequently identified by PCR. One alternative method for identifying the ChIP-isolated DNA binding regions is by hybridization to DNA microarrays (ChIP-chip) [54,55].

WGas can be used for an unbiased, genome-wide identification survey of in vivo transcription factor binding sites by chromatin immunoprecipitation coupled with array hybridization [55]. The high reproducibility among arrays, unbiased and complete genomic coverage, and multiple,
potentially overlapping, probes representing transcription factor binding regions are significant advantages of tiling arrays over other array designs. Indeed, at a high probe tiling resolution, multiple overlapping probes may contain the actual transcription factor binding motif and thus enable a fine mapping of the binding site to a resolution of less than 25 bp. Although it may not yet be practical in terms of cost and the number of chips required for some organisms, when used for ChIP-chip studies high-density oligonucleotide arrays coupled with rigorous statistical analysis methods can provide high-resolution binding site location data with high sensitivity and reasonably low false-positive rates.

Microarrays containing genomic PCR products have been used for ChIP-chip studies in yeast [56-62] and in mammalian cells [56,63-66]. However, many of the ChIP-chip studies in mammalian cells have analyzed only selected predicted promoter regions and do not interrogate the entire genome. Such arrays cannot address the possibility that transcription factors might bind at other locations. For example, it has been shown in yeast that forkhead transcription factors can associate with the genomic coding regions and function to coordinate transcription elongation and transcript processing [67].

Recently several unbiased chromosome-wide transcription factor studies in mammalian cells have been undertaken. A somewhat surprising finding from this work has been that a large fraction of in vivo binding sites were outside of the predicted promoter regions of genes [8,63,68]. It was found that the binding locations of NF-κB [63]; cMyc, Sp1, and p53 [8]; and CREB [68] were located within both coding and noncoding regions. More binding sites were found than expected, and only a relatively small fraction of the sites occurred in regions that would typically be considered “promoters”—near the 5’ ends of protein-coding genes. The binding sites frequently occurred in introns, coding regions, or unannotated transcriptionally active regions or near the 3’ ends of genes. Collectively, these results indicate that transcription factor binding is not restricted to canonical annotated upstream promoter regions and may herald a paradigm shift for our understanding of transcriptional regulation. Unbiased genome tiling arrays were key to these discoveries.

Methylome analysis

Chromatin modifications associated with regulation of gene expression include methylation of cytosines as well as the acetylation, methylation, phosphorylation, etc., of histones [69]. In eukaryotes, cytosine methylation has evolved into a mechanism that allows dividing cells to inherit states of gene activity stably, and in most cases DNA methylation causes gene silencing. The most conserved role for DNA methylation in eukaryotes is in the silencing of repetitive elements in the genome, primarily transposons and retroviruses [70,71]. DNA methylation is also important in cancer biology, as tumors often show both hypomethylation and hypermethylation of specific tumor suppressor genes. DNA methylation is generally thought to have initially evolved as a defense mechanism against “foreign” DNA sequences that pose a threat to genome stability (such as transposable elements and retroviruses) [71,72] and later assumed other cellular functions, such as developmental regulation of gene expression. In addition to its role in transposon silencing, DNA methylation is involved in a myriad of epigenetic regulatory processes found throughout the eukaryotes. In mammals, cytosine methylation plays a key role in two gene regulatory systems that involve allele-specific methylation, mammalian parental imprinting [73], and X-chromosome inactivation [74]. DNA methylation is also involved in cancer biology [75,76]. Tumors consistently show abnormal patterns of methylation including genome-wide demethylation and inappropriate hypermethylation of particular genes. Methylation of genomic regions is significant, as heavy cytosine methylation is almost always associated with histone modifications (e.g., histone H3 lysine 9 methylation), chromatin condensation, and repression of gene expression. Despite the importance of DNA methylation, and despite decades of phenomenological descriptions of epigenetic regulatory systems, genetic studies of the mechanisms of DNA methylation control are still in their infancy. In part this is because, even though methylation is present in most eukaryotes, it has been curiously lost in some of the most well studied model organisms including yeast and Caenorhabditis elegans. Drosophila has a very small amount of methylation, but has lost most of the methylation machinery found in other eukaryotes [77].

Bisulfite genomic sequencing is the best technique for mapping DNA methylation at single-base resolution, but this technique has not yet been applied to whole-genome approaches. Sodium bisulfite treatment of DNA deaminates cytosine to uracil, but 5’-methylcytosine is protected [78]. Thus DNA methylation is detected as cytosine residues that survive bisulfite treatment and can be assayed by PCR amplification and DNA sequencing or by restriction enzyme digestion followed by PCR or DNA hybridization. Sodium bisulfite conversion can be utilized for microarray analysis because unmethylated DNA that is treated with bisulfite contains uracil in place of cytosine and will hybridize relatively poorly to microarray oligonucleotides that contain guanines. Methylated cytosines in DNA sequences that cannot be changed by bisulfite treatment will retain their ability to hybridize to these oligos (A. Sundaresan and J.R. Ecker, unpublished). The bisulfite microarray method has several advantages for assaying DNA methylation. Because there is no need to fractionate the DNA, every nucleotide of the genome is inspected, and since hybridization to a particular oligo is affected by the methylation pattern present in that 25-mer sequence, regions of methylated DNA can be accurately defined with single-probe resolution. Moreover, the effect of sodium bisulfite treatment on the hybridization of a given microarray element can be
predicted, adding to the usefulness of this method. However, the major limitation of the bisulfite method is that some regions of lightly methylated DNA may not be able to hybridize to array oligos after bisulfite conversion even if the region is partially methylated. Bisulfite conversion coupled with microarray technology has been used to detect methylated DNA regions (http://innovation.swmed.edu/research/instrumentation/higher_pages/inst_doc_about.html). In this work, genomic DNA fragments are hybridized to the oligonucleotide resequencing arrays following bisulfite treatment. In this way the methylation status of several promoter regions was analyzed at 1-base resolution.

Another potentially useful approach to methylome analysis would involve using methylation-sensitive restriction enzymes. This method exploits the specificity of bacterial restriction enzymes to distinguish methylated and unmethylated DNA sequences. Endonucleolytic cleavage of DNA can destroy hybridization to a 25-mer oligonucleotide, particularly if the restriction enzyme site lies close to the middle of the element. Therefore, oligonucleotides on the microarray that contain restriction sites will show reduced hybridization if the site is cleaved, allowing methylated DNA sequences to be identified. Hybridization of this DNA to a tiling microarray and comparison with labeled fragments generated from undigested DNA will show a reduction of hybridization wherever a site has been cleaved in the middle of an oligonucleotide element. This technique is feasible only with oligonucleotide arrays containing relatively short (i.e., 25-mer) probes because long oligos or larger PCR products will retain hybridization after restriction digestion. An elaboration of this method involves restriction enzyme digestion coupled with size selection to discriminate between methylated and unmethylated DNAs [79] and will detect regions of light DNA methylation more effectively than restriction digest without size fractionation. In this study sucrose gradient sizing and labeling of small fragments of DNA that had been digested with a methyl-sensitive restriction enzyme were used and the products were then hybridized to an array that contained 384 PCR fragments across the genome [79].

Additional approaches for methylome analysis involve the identification of methylated DNA by immunoprecipitation or affinity chromatography, followed by hybridization of the purified genomic fragments to microarrays. Methylated DNAs can be isolated by immunoprecipitation of fragmented DNAs with anti-5-methylcytosine antibodies, as has been shown previously [80], and then the purified products could be labeled and hybridized to tiling arrays. Alternatively, methylated DNA could be purified by affinity chromatography. Methyl-binding domains are found in a family of eukaryotic chromatin proteins and specifically recognize methylcytosine. For example, MeCP2 will bind if only a single methylated CG is present [81], and it has been shown that methyl-binding-domain (MBD) agarose chromatography fractionates DNA according to its degree of CG methylation [82]. This approach was recently used to survey the methylated component of the Neurospora crassa genome, by large-scale sequencing of fragments purified by MBD agarose chromatography [83]. Thus, affinity-purified methylated gDNA fractions could be hybridized to tiling microarrays to identify unmethylated, lightly methylated, and heavily methylated portions of the genome. Both immunoprecipitation and affinity chromatography are likely to have extremely high sensitivity for detecting lightly methylated DNA sequences; however, one limitation of these approaches is that the purified fragments will be relatively long (~200–500 bp). Although these methods would be very useful in defining all of the methylated regions in the genome, they will not precisely identify the DNA methylation at specific cytosine residues.

Methods such as those described above have not yet been applied to whole-genome tiling arrays to survey comprehensively the methylome at high resolution. In principle, oligonucleotide tiling arrays that cover the entire genome will allow a truly genome-wide DNA analysis of DNA methylation. Because genome tiling arrays can be synthesized to contain relatively short (25-nucleotide) oligonucleotides, they can potentially identify sites of DNA methylation with unparalleled precision, in some cases with single-nucleotide resolution. Because tiling arrays are universal platforms they are ideal for assessing the correlation of a variety of types of genomic information. WGAs can be used for detecting correlations between DNA methylation and global transcriptome mapping and between patterns of DNA methylation with sense transcripts, antisense gene transcripts, or noncoding transcripts on a whole-genome scale using wild-type tissues, cancer cells, or specific genetic mutant backgrounds (i.e., mutations in specific methylases, acetyltransferases, DNA binding proteins, etc.).

Polymorphism analysis and genome resequencing

WGAs can be used for several related approaches aimed at comparing genomic DNA sequences to a reference sequence and thus identifying genome-wide sequence polymorphisms in a single array hybridization experiment. Comparative genome hybridization (CGH), single-feature polymorphism (SFP) detection and genotyping, and tiling array-based genome resequencing essentially represent whole-genome resequencing at low, medium, and high resolution, respectively. CGH has been used mainly to identify relatively large insertions and deletions in cancer cell lines by hybridization of labeled DNA from different cell lines either to arrays of BAC clones tiled to cover the whole genome [84] or to cDNA expression arrays [85]. More recently, tiling DNA arrays containing BAC clones representing the entire human genome [86] and oligonucleotide arrays [87] have been used for CGH. Clearly, WGAs can be useful for high-resolution CGH.
Another potential application of WGAs is for the characterization and detection of DNA sequence variations in individuals or populations compared to a reference genome sequence. Identifying and cataloging such polymorphisms is an important step in positional cloning and candidate gene strategies for the identification of genes associated with disease or interesting phenotypes. Array hybridization techniques have been used successfully to study SFPs, which are identified when specific 25-mer oligonucleotide features exhibit significantly different hybridization signals when different strains within a species are compared [88,89]. This approach identifies SFPs at a resolution of 25 bp, but does not provide the actual sequence change of the polymorphism. In recent studies, standard Affymetrix expression arrays have been used to identify thousands of SFPs among different strains of *Arabidopsis* [89] (Borevitz and Ecker, unpublished; http://naturalvariation.org/sfp). It is obvious that this approach would benefit tremendously from the use of whole-genome tiling arrays in place of the limited number of probes available on existing expression arrays. Not only would WGAs enable the identification of many more SFPs, but a WGA design using partially overlapping probes may yield a potential SFP resolution of less than 25 bp.

The highest resolution and potentially most powerful tiling WGA designs are intended for complete resequencing of a genome relative to the reference sequence by hybridization [90]. Such array-based DNA sequencing is dependent on the prior availability of a finished quality reference sequence and is thus an alternative method for resequencing the same genome. This application requires a specialized ultra-high-resolution (1 bp) high-density array design [91,92]. Resequencing arrays are designed using a set of tiled, overlapping oligonucleotide probes. For each base being interrogated there are four probes, one representing the reference sequence and three that vary the central base with one of the other three nucleotides. A mismatch (SNP) at the middle base will reduce or prevent binding of the labeled target and thus result in a reduced hybridization signal (Fig. 2D). Interrogation of the sequence of both strands of a chromosome of length \(N\) requires \(8N\) oligonucleotide probes. Using this approach an entire genome can be resequenced in a single hybridization experiment [93]. Currently one technical challenge for resequencing arrays is the amplification and labeling of the target, which typically involve long-range PCR. However, for organisms with a small genome direct labeling of genomic DNA may be possible, bypassing the need for PCR amplification prior to hybridization. Because resequencing arrays can compare a new sequence only to the reference sequence on which they are based they will not be as useful for identifying novel sequences or sequence rearrangements that may occur in the experimental sample. The large number of probes required for complete resequencing of a typical eukaryotic genome may make array-based resequencing seem impractical. However, given the recent increases in feature densities, it would require only ~300 chips to resequence the entire ~125-Mb *Arabidopsis* genome using oligonucleotide arrays. Moreover, if current trends in feature size continue to follow Moore’s Law, then this technology may soon allow for routine resequencing of much larger genomes (Fig. 4).

**Tiling array design considerations**

While tiling the entire genome with oligonucleotide probes provides an unbiased view of the genome, there are specific design and analysis problems that arise. For expression arrays, probes are carefully selected and represent only a small subset of the possible probes within exons [94,95]. Probes are chosen so that they all hybridize with similar efficiencies at a given temperature and are designed to be nonpalindromic to prevent self-hybridization. Expression chip probes are generally selected to minimize cross-hybridization with other known parts of the transcriptome. In addition, gene expression probe sets are selected to provide a linear signal response with respect to target concentration. When tiling the entire genome, these parameters are impossible to maintain.

The simplest approach to designing a whole-genome tiling array is to start at the beginning of each chromosome and cover the entire sequence with 25-mer probes tiled end to end and, for higher resolution, doing this multiple times with an offset between tile phases [11]. However, many of these
the probes chosen with this simple strategy may be problematic; some would be duplicated, some would have weak hybridization affinity, and some would hybridize so strongly that even nonspecific background hybridization would generate a saturated signal. Also, some probes may have to be eliminated because of sequence-specific fabrication constraints. One alternative is to modify the design from a perfect end-to-end tiling with no overlap to a design allowing for varying amounts of overlap, or gaps, between probes.

Standard genomic repeat masking (i.e., RepeatMasker: http://www.repeatmasker.org/) prior to probe selection is currently a common first step. Probes that are identical repeats at the individual probe sequence level may also be eliminated from consideration because some significant problems may arise for probes that occur many times in the genome. Such probes are uninformative since the precise genomic location of the signal from that probe will be uncertain. Moreover, extreme signals from repeated probes representing highly transcribed regions may adversely affect background correction and normalization routines and thus present data analysis problems. Further filtering of probes could eliminate probes predicted to have such a low binding affinity as to be essentially useless and also those probes with such high affinity that they would be uninformative due to saturating cross-hybridization and may also introduce problems for background correction and normalization. Models of DNA duplex and RNA–DNA heteroduplex hybridization for both perfect-match and single-base mismatches in solution based on oligonucleotide hybridization theory have been developed to estimate the binding affinity and specificity of oligonucleotide probes. Nearest-neighbor models approximate the binding affinity of a probe based on the affinity of each base pair depending on the sequence context [96–103]. Because the probes on arrays are tethered by a linker to the chip surface and are not really in solution, Langmuir surface adsorption models [104–106] have also been developed. An alternative empirically derived composite physical/stochastic model has been proposed as a complement to the physical models [107].

Probe affinity modeling and determinations of probe specificity on a whole-genome level can be used to screen candidate oligos and eliminate those that are likely to be problematic from the chip design. The thresholds chosen for exclusion of some probes based on affinity considerations must be loose enough to avoid introducing significant bias into the design and thus leave large portions of the genome uninterrogated, but must simultaneously be restrictive enough to introduce savings in terms of chip fabrication costs or data analysis. Alternatively, probe affinity predictions can be used to adjust or correct hybridization signals during data processing. Another consideration is whether only perfect-match probes or perfect-match/mismatch (PM/MM) probe pairs should be used. Using PM/MM probe pairs may have some advantages in correcting for nonuniform probe affinities, but it cuts in half the amount of genomic sequence that can be represented on a single chip. Some newer analysis methods do not consider the MM intensity (e.g. RMA [108], GCRMA [109], fitPLM [110]); however, the relative strengths and weaknesses of these different approaches remain an area of active debate [107,108].

Challenges and future directions

While whole-genome tiling arrays are already a powerful tool for genomic analysis, at present there remain significant practical limitations imposed by the number of unique probe features that can be synthesized on a single chip and thus how many chips are required to cover an entire genome. For example, a tiling array design representing human chromosomes 21 and 22 and interrogating ~35 Mb of nonrepetitive sequence with probes spaced about every 35 bases on average required 3 chips [9]. However, a more recent design to tile about 30% of the human genome (~490 Mb) with overlapping 25-mer probes spaced every 5 bases requires 98 chips/experiment (T. Gingeras, personal communication). For smaller genomes such as Arabidopsis, a set of only 12 chips was required to tile both strands of virtually the entire ~125-Mb genome at 25-base resolution, and a complete Arabidopsis tiling array design at ~8-base resolution required only 36 chips [11].

While full genome coverage of a large eukaryotic genome (~3 Gb) can theoretically be obtained, many more chips would be required. Thus, from a cost perspective, such experiments would be impractical. Even annotation-directed array designs that do not attempt to represent the entire human genome sequence require many chips. For example, a set of 50 ink-jet-synthesized arrays was needed to probe about 443,000 Genscan predicted exons in the human genome, with only a single oligo per exon [13], and a study of 52 different samples using ink-jet arrays containing ~125,000 splice-junction probes for only 10,000 genes used 260 arrays (single oligo per exon) [12]. However, considering that the increasing feature density of successive generations of oligonucleotide arrays appears to follow a type of Moore’s Law (Fig. 4), future generations of tiling arrays can be expected to provide much higher feature densities. Future improvement on WGA technology may even allow for tiling an entire hundred-megabase- to gigabase-scale genome on a single array, making it practical to use whole-genome arrays as a universal platform for a variety of unbiased genomic studies. Such improvements in feature density will make it practical to design and pursue experiments with numerous different samples and enable sufficient biological replication using independent biological samples instead of technical replicates of the same biological material. Furthermore, the ability to tile the genome at a higher resolution (i.e., more overlap between the oligos) may make it possible to identify precisely the boundaries of exons and introns, to identify miniexons [111] that are shorter than the probes
themselves (i.e., <25 bp), and to define individual transcription factor binding sites in ChIP-chip studies [8].

There is also considerable room for improvement in other areas. The utility of tiling arrays for empirical annotation can be expected to improve if transcription populations representing a large variety of tissues, cell types, genotypes, and treatments are assayed, thereby increasing the likelihood of identifying low-abundance and/or rare transcripts. The power of tiling arrays as a gene discovery tool will improve with increased sensitivity that may be achieved through experimental replication and will benefit from improved analytical techniques that account for the affinity characteristics of individual oligonucleotide probes. Improved methods for capturing low-abundance messages and generating labeled targets representing full-length transcripts [40] will be needed. Since an intact transcript is not required, subtractive hybridization of mRNA populations may be useful. Methods for isolating and labeling atypical RNAs will be necessary to characterize the full transcriptome. A growing body of evidence increasingly points to the existence of a “hidden transcriptome,” which until very recently was considered “background” transcriptional activity. Recent studies, however, are revealing a widely expanded role for noncoding RNAs that was previously unimaginable [19,112,113]. Both genetic studies and the use of tiling array technology may reveal that the transcriptional “noise” in the genome may actually turn out to be music that the genome has been playing but we were unable to appreciate fully.

Additional challenges will include cataloging and analyzing the enormous amount of data generated even from a single whole-genome tiling array experiment. User-friendly graphical interfaces such as ChipViewer (Fig. 3; H. Chen and J. Ecker, unpublished) for the visualization and integration of WGA data with other data and annotations must be further developed. Moreover computational approaches are need for integrating these data with other genome annotations such as DNA methylation and protein binding sites. New algorithms and statistical methods for data analysis will be needed to extract meaningful results from these massive datasets. For example, one of the goals of empirical transcriptome mapping using tiling arrays is to infer transcript structures for genes that may be incorrectly predicted by gene prediction algorithms or missed altogether. At present there are no standard methods or analytical tools for scoring tiling array probe intensities to infer transcriptional activity or predict transcript structures. One approach, using a sliding window method to merge the discontinuous hybridization signals of probes representing discrete exons together to predict transcript structures, has been developed for this purpose [10]. Alternatively, Bayesian models of probe responses that are trained using probe signals for known full-length transcripts may be used to predict likely transcript structures for hypothetical genes or novel transcripts in intergenic regions (Mockler and Ecker, unpublished). However, problems such as nonuniform probe responses due to varying probe affinities are likely to confound such ad hoc solutions.

Another key challenge involves the validation of possible “paradigm-shifting” discoveries made through experiments using tiling arrays. Comparison of results using different model organisms (e.g., bacteria, yeast, flies, worms, mammals, and plants) and data obtained using experimental approaches (e.g., MPSS [114–117], SAGE [118,119]), chip designs, and analytical methods will be needed. At present, discoveries made using tiling array technology require confirmation with more conventional methods such as directed RT-PCR, cloning, and sequencing. However, rapid-prototyping technologies for chip design and manufacture such as so-called “maskless” photolithography technology [5–7] may make it possible to validate quickly findings made using one chip design with a completely new chip design.

Note added in proof


Acknowledgments

We thank Jennifer L. Nemhauser, Justin Borevitz, and Todd P. Michael for critically reading the manuscript; Pam Surko for helpful discussions and valuable insights; Joanne Chory for advice and support; Huaming Chen for the data and analysis shown in Fig. 3; T. Gingeras for sharing unpublished data; and an anonymous reviewer for making constructive comments. T.C.M. is an NIH Postdoctoral Fellow (F32 GM69090). This work was supported by grants from the National Science Foundation 2010 Program and the National Institutes of Health ENCODE Program to J.R.E.

References


[110] B.M. Bolstad, Low level analysis of high-density oligonucleotide array data: background, normalization and summarization, Biostatistics, Univ. of California, Berkeley, 2004.


